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(54)Porous tissue scaffoldings for the repair or regeneration of tissue

The present patent describes a three-dimensional interconnected open cell porous foams that have a gradient in composition and/or microstructure through one or more directions. These foams can be made from a blend of absorbable and biocompatible polymers that are formed into foams having a compositional gradient transitioning from predominately one polymeric material to predominately a second polymeric material. These gradient foams are particularly well suited to tissue engineering applications and can be designed to mimic tissue transition or interface zones.

Description

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Field of the Invention

[0001] The present invention relates generally to the field of tissue repair and regeneration. More particularly the present invention relates to provise biocompatible bloabsorbable foams that have a gradient in composition and/or microstructure that serve as a template for tissue regeneration, repair or automentation.

Background of the Invention

[0002] Open cell porous biocompatible foams have been recognized to have significant potential for use in the repair and regeneration of tissue. Early efforts in tissue repair focused on the use of amorphous biocompatible foam as porous plugs to fill voids in bone. Brekke, et al. (U.S. Patent No. 4,186,448) described the use of porous mesh plugs composed of polyhydroxy acid polymers such as polylactide for healing bone voids. Several attempts have been made in the recent past to make TE scaffolds using different methods for example U.S. Patents 5,522,895 (Mikos) and 5,514,378 (Mikos. et al.) using leachables; U.S. Patents 5,755,792 (Brekke) and 5,133,755 (Brekke) using vacuum foaming techniques; U.S. Patents 5,716,413 (Walter, et al.) and 5,607,474 (Athanasiou, et al.) using precipitated polymer gel masses; U.S. Patents 5,686,091 (Leong, et al.) and 5,677,355 (Shalaby, et al.) using polymer melts with fugitive compounds that sublimate at temperatures greater than room temperature; and U.S. Patents 5,770,193 (Vacanti, et al.) 5,769,899 (Schwartz, et al.) and 5,711,960 (Shikinami) using textile-based fibrous scaffolds. Hinsch et al. (EPA 274,898) described a porous open cell foam of polyhydroxy acids with pore sizes from about 10 to about 200 µm for the in-growth of blood vessels and cells. The foam described by Hincsh could also be reinforced with fibers, yarns, braids, knitted fabrics, scrims and the like. Hincsh's work also described the use of a variety of polyhydroxy acid polymers and copolymers such as poly-L-lactide, poly-DL-lactide, polyglycolide, and polydioxanone. The Hincsh foams had the advantage of having regular pore sizes and shapes that could be controlled by the processing conditions, solvents selected, and the additives

[0003] However, the above techniques have limitations in producing a scaffold with a gradient structure. Most of the scaffolds are isotropic in form and function and lack the anisotropic features of natural tissues.

[0004] Further, it is the limitation of prior art to make 30 scatfolds that have the ability to control the spatial distribution of various pore shapes. The process that is described to fabricate the microstructure controlled foams is a low temperature process that offers many advantages over other conventional techniques. For example the process allows the incorporation of thermally sensitive compounds like proteins, drugs and other additives with the thermally and hydrolytically unstable absorbable polymers.

[0005] Althanasiou et al. (U.S. Patent No. 5607.474) have more recently proposed using a two layer foam device for repairing osteochondral defects at a location where two disminst types of tissue are present. The Althanasiou device is composed of a first and second layer, prepared in past separately, and pined together at a subsequent step. Each of the scatflotd layers is designed to have stiffness and compressibility corresponding to the respective cartilage and bone test. Since cartillage and bone test layers in the both its approach is an attempt to more clearly mimic the structure of the human body. However, the interface between the cartilage and bone in the human body is not a discretel junction of two dissimiter materials with an abrupt change in anatomical features and/or the mechanical properties. The cartilage cells have distinctly different cell morphology and orientation depending on the location of the cartilage cell in relation to the underlying bone structure. The difference in cartilage cell morphology and orientation depending on the location of the cartilage cell in relation to the underlying bone structure. The difference in cartilage cell morphology and privately the cartilage cell morphology and privately in the topic of the cartilage cell morphology and privately the cartilage cell in relation to the underlying bone cartilage cell morphology and privately the cartilage cell morphology

[0006] Another approach to make three-dimensional laminated foams is proposed by Mikos et al. (U.S. Patent No. 5,514,378). In this technique which is quite cumbersome, a porcus membrane is first prepared by drying a polymer solution containing leachable salt crystals. A three-dimensional structure is then obtained by laminating several membranes together, which are cut to a contour drawing of the desired shape.

[0007] One of the major weaknesses of the prior art regarding three-dimensional porous scaffolds used for the regeneration of biological tissue like cartilage is that their microstructure is random. These scaffolds, unlike natural tissue, or to not vary in morphology or structure. Further, current scaffolds do not provide adequate nutrient and fluid transport for many applications. Finally, the laminated structures are not completely integrated and subjected to delamination under in vivo conditions.

[0008] Therefore, it is an object of the present invention to provide a biocompatible, bioabsorbable foam that provides a continuous transitional gradient of morphological, structural and/or materials. Further, it is preferred that foams used in tissue engineering have a structure that provides organization at the microstructure level that provides a template that facilitates cellular invasion, proliferation and differentiation that will ultimately result in regeneration of functional

tissue.

Summary of Invention

[0009] The present invention provides a biocompatible gradient form that has a substantially continuous transition in at least one characteristic selected from the group consisting of composition, stiffness, flexibility, bioebsorption rate pore architecture and/or microstructure. This gradient form can be made from a blend of absorbable polyment at form compositional gradient transitions from one polymeric material to a second polymeric material, in situations where a single characterial composition is sufficient for the application, the invention provides a biocompatible clear that may have microstructural variations in the structure across one or more dimensions that may minic the anatomical features of the tissue (e.g. cartillage, skip, bone etc.).

[Ootig] The present invention further provides biocompatible foam having interconnecting pores and channels to facilitate the transport of nutrients and/or invasion of cells into the scallold. These biocompatible foams are especially well adapted for facilitating the ingrowth of tissue as is described in Example 7.

[0011] In yet another embodiment of the present invention biocompatible foams having interconnecting pores formed from a composition containing in the range of from about 30 weight percent to about 99 weight £ caprolactone repeating, units are disclosed. These biocompatible foams are especially well adapted for facilitating the growth of osteoblasts as is described in Example 6.

[0012] The present invention also provides a method for the repair or regeneration of tissue contacting a first tissue with a gradient foam at a location on the foam that has appropriate properties to facilitate the growth of said tissue. The concept of a continuous transition in physical properties, chemical composition and/or microstructural features in the porcus scaffold (loam) can facilitate the growth or regeneration of tissue. These foam structures are particularly settly for the generation of tissue ignations between two or more different types of tissues. For a multi-cellular eystern in the simplest case, one cell type could be present on one side of the scaffold and a second cell type on the other cise of the scaffold. Examples of such regeneration can be (a) skirt, with throbotasts on one side to regenerate domins, and feratinocytes on the other to regenerate epiderms; (b) vascular grafts: with an endothelial layer on the inside of

Brief Description of Figures

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[0013] Figure 1 is a scanning electron micrograph of the cross section of a random microstructure foam made from 5% solution of 35/65 e-caprolactone-co-glycolide copolymer.

[0014] Figure 2 is a scanning electron micrograph of the cross section of a foam with vertical open channels made from 10% solution of 35/65 e-caprolactone-co-glycolide copolymer.

[0015] Figure 3 is a scanning electron micrograph of the cross section of a foam with architectural gradient made from 10% solution of 35/65 e-caprolactone-co-glycotide copolymer.

[0016] Figure 4 is a scanning electron micrograph of the cross section of a gradient foam made from a 50/50 blend of 40/60 ε-caprolactone-co-(L)lactide copolymer and 35/65 ε-caprolactone-co-glycolide copolymer.

[0017] Figure 5 is a scanning electron micrograph of a cross section of the top portion of a gradient foam made from a 50/50 blend of 40/50 e-caprolactone-co-(L)lactide copolymer and 35/65 e-caprolactone-co-glycolide copolymer.

[0018] Figure 6 is a scanning electron micrograph of a cross section of the bottom portion of a gradient foam made from a 50/50 bend of 40/50 e-caprolactone-co-(L)lacitie copolymer and \$565 e-caprolactone-co-glycolide copolymer.

[0019] Figure 7 is a graphical presentation of cell culture data, 77, 78 and 70.

[0020] Figure 8 is an anatomical sketch of cartilage tissue.

[0021] Figure 9A, 9B, and 9C are scanning electron micrographs of a 0.5 mm foam made from a 50/50 blend of a 35/65 e-caprolactione-or-glycolide copolymer and a 40/60 e-caprolactione-or-(L)actide copolymer with architecture suitable for use as a skin scallodid. Figure 9A shows the porosity of the surface of the scalfold that preferably would lace the wound bed. Figure 9B shows the porosity of the surface of the scalfolding that would preferably face away from the wound bed. Figure 9C shows a cross section of the scalfold with channels running through the thickness of

[0022] Figure 10 is a dark field 40X photomicrograph of a trichrome stained sample illustrating the cellular invasion of the foam shown in Figure 9, eight days after implantation in a swine model.

[0023] Figure 11 is a 100X composite photomicrograph of a trichrome stained sample illustrating the cellular invasion of the feam shown in Figure 9 which also contained PDGF, eight days after implantation in a swine model.

Detailed Description of the Invention

[0024] This invention describes porous bloabsorbable polymer foams that have novel microstructures. The features

of such foams can be controlled to suit a desired application by choosing the appropriate conditions to form the foam during lyophilization. These features in absorbable polymers have distinct advantages over the prior ant where the scaffolds are lysically isotropic or random structures. However, it is preferred that foams used in tissue engineering (i. e. repair or regeneration) have a structure that provides organization at the microstructural level that provides a template that facilitates cellular organization and regeneration of tissue that has the antentical, biomechanical, and biochemical features of normal tissues. These foams can be used to repair or regenerate tissue (including organs) in animals such as domestic animals, primates and humans.

[0025] The features of such foams can be controlled to suit desired application by selecting the appropriate conditions for tyophilization to obtain one or more of the following properties: (1) interconnecting pores of sizes ranging from about 10 to about 200 µm (or greater) that provide pathways for cellular ingrowth and nutrient diffusion; (2) a variety of porosities ranging from about 20% to about 98% and preferably ranging from about 80% to about 95%; (3)gradient in the pore size across one direction for preferential cell culturing; (4) channels that run through the foam for improved cell invasion, vascularization and nutrient diffusion; (5) micro-patterning of pores on the surface for cellular organization: (6) tailorability of pore shape and/or orientation (e.g. substantially spherical, ellipsoidal, columnar); (7) anisotropic mechanical properties; (8) composite foams with a polymer composition gradient to elicit or take advantage of different cell response to different materials; (9) blends of different polymer compositions to create structures that have portions that will break down at different rates; (10) foams co-lyophilized or coated with pharmaceutically active compounds including but not limited to biological factors such as RGD's, growth tactors (PDGF, TGF-β, VEGF, BMP, FGF etc.) and the like; (11) ability to make 3 dimensional shapes and devices with preferred microstructures; and (12) lycobilization with other parts or medical devices to provide a composite structure. These controlled features in absorbable polymers have distinct advantages over the prior art where the scaffolds are typically isotropic or random structures with no preferred morphology at the pore level. However, it is preferred that foams used in tissue scaffolds have a structure that provides organization at the microstructure level and provides a template that facilitates cellular organization. ization that may mimic natural tissue. The cells will adhere, proliferate and differentiate along and through the contours of the structure. This will ultimately result in a cultured tissue that may mimic the anatomical features of real tissues to a large extent.

[0026] For example, as shown in Figure 3 the orientation of the major axis of the pores may be changed from being in the same plane as the foam to being oriented perpendicular to the plane of the foam. As can be sent from Figure 3 the pore size can be varied from a small pore size generally between about 30 µm and about 50 µm to a larger size of from about 100 µm to about 200 µm in porous gradient foams, Ideally the foam structure could be created to facilitate the repair or regeneration of human tissue junctions such as the cartilage to bone junction present in joints. This foam would progress from a small (i.e. about 30 µm to about 150 µm in diameter) ound pore to larger column-tike pores (i.e. about 30 µm to about 400 µm in diameter, preferably about 150 µm to about 400 µm in diameter, in most cases with a length to diameter ratio of at least 22. Feams with channels are illustrated in Figure 2 and Figure 3. The channels formed by this process generally begin on one surface of the foam and may traverse the thickness of the foam. The channels length is generally at least two times the average pore diameter. Amonels for most perferably at least eight times the average pore diameter. Channels for most perferably at least two times the average pore diameter. Amonels for most perferably at least two times the average pore diameter. Channels or most perferably at least 20 or diameter. Amonels for most perferably at least 20 or diameter. Amonels or most perferably at least 20 or diameter. Amonels or most perferably at least 20 or diameter and preferably at least 20 or diameter. Amonels are all seast one time the size of the average pore diameter and preferably at least 20 or diameter. Or diameter of course will be selected based on the desired functionality of the channel such as cellular invasion, nutrient diffusion or as an avenue for vascularization.

[0027] There are a number of biological tissues that demonstrate gradient architectures. Examples of tissues where a gradient scalloid could be used, include, but are not limited to: bone, spine disc, articular cartilage, meniscus, fibrocartilage, tendons, ligaments, dura, skin, vascular grafts, nerves, liver, and pencreas. The examples below only highlight a few tissues where gradient scalloids could be used. The design of tissue engineered scalloids to facilitate development of these organ structures would benefit greatly from the ability to process or create a gradient architecture in the scalfold.

Cartilage

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[0028] Articular cartilage covers the ends of all bones that form articulating joints in humans and animals. The cartilage acts in the joint as a mechanism for force distribution and as a bearing surface between different bones. Without articular cartilage, stoses concentration and friction would occur to the dogree that the joint would not permit ease of motion. Loss of the articular cartilage usually leads to painful arthritis and decreased joint motion. A schematic showing the morphological features of a healthy cartilage is shown in Figure 8.

[0029] Articular cartilage is an excellent example of a naturally occurring gradient structure. Articular cartilage is composed of four different zones that funded the superficial or tangenistal zone within the first 10-20% of the structure. (this includes the articular surface), the middle zone which is 40-60% of the middle structure, and the deep zone than the surface of the structure.

is adjacent to the tide mark, and a transition zone between the bore and cartilage that is composed of calcified cartilage. Subchondral bone is located adjacent to the tide mark and this transitions into cancellous bone. In the superficial or tangential zone, the collegen tibrils are parallel to the surface. The fibers are oriented to resist shere forces generated during normal joint articulation. The middle zone has a randomly arranged organization of much larger diameter collegen fibers. Finally, in the deep zone there are larger collegen fiber bundles, which are perpendicular to the surface, and they insert into the calcilied cartilage. The cells are specified and tend to arrange themselves in a columnar manner. The calcilided cartilage zone has smaller cells with relatively tittle evologism.

manner. The calcified cartilage zone has smaller cells with relatively little cytoplasm. [0030] A preferred embodiment of this invention would be to generate a gradient foam structure that could act as a template for multiple distinct zones. These foam structures could be fabricated in a variety of shapes to regenerate or repair osteochondrial defects and cartilage. One potential foam structure would be cylindrical in shape with an approximale dimensions of 10mm in diameter and 10 mm in depth. The top surface is would be approximately 1 mm thick and would be a low porosity layer to control the fluid permeability. By adopting a suitable processing method the surface porosity of the foam could be controlled. The porosity of this skin like surface can be varied from completely impervious to completely porous. Fluid permeability would be controlled by surface porosity. Below such a skin the structure would consist of three zones. An upper porous zone which lies adjacent to cartilage tissue, a lower porous zone which lies adjacent to bone tissue, and a transition zone between the upper and lower porous zones. For articular cartilage, it is currently preferred that the stiffness (modulus) of the upper and lower porous layers at the time of implantation be at least as stiff, as the corresponding adjacent tissue. In such a case the porous layers will be able to support the environmental loading and thereby protect the invading cells until they have differentiated and consolidated into tissue that is capable of sustaining load. For example the porous structure used for the superlicial tangential zone could have elongated pores and the orientation of the structure could be parallel to the surface of the host cartilage. However, the deep zone may have a porosity of about 80 to about 95 % with pores that are of the order of 100 µm (about 80 µm to about 120 μm). It is expected that chondrocytes will invade this zone. Below this, would be a zone with larger pores (about 100 μm to about 200 μm) and a porosity in the range of about 50 to about 20 %. Such 100 μm to about 200 μm porous foam would have a structure such that the struts or walls of the pores are larger and vertical to the load, similar to the naturally occurring structure and to bear the loads. Finally, at the bottom of this structure there is a need for larger peres (about 150 µm to about 300 µm) with higher stiffness to be structurally compatible with cancellous bone. The foam in this section could be reinforced with ceramic particles or fibers made up of calcium phosphates and the like. [0031] Recent data generated in our laboratories support the hypothesis that cell invasion can be controlled by pore size. In these studies, a scaffold made of 95/5 mole percent poly(L)lactide-co-ε-caprolactone) with an approximate pore size of about 80 µm had chondrocyte invasion of about 30 cells/mm² of the scaffold (under static conditions). Scaffolds made of 40/60 mole percent poly(ε-caprolactone-co-(L)lactide) with a larger approximate pore size of about 100 µm had a statistically significantly greater cellular invasion of 50 cells/mm² (under static conditions). In both cases the cells were bovine chondrocytes. A very simple gradient structure with a variation of pore sized from about 80 µm to about 150 µm would provide a structure where chondrocytes would more easily invade the area with larger pores. The area with smaller pores would be void of chondrocytes or would be filled with a second cell types (e.g., fibroblasts). [9032] In a compositionally gradient foam a blend of two or more elastomeric copolymers or in combination with high modulus semi-crystalline polymers along with additives such as growth factors or particulates can be chosen such that first a desired pore gradient is developed with a preferred spatial organization of the additives. Then using a variety of the approaches referred to in the preferred methods of making gradient foams, a compositional gradient can be superimposed primarily due to the differences in the polymer-solvent phase separation behavior of each system. Such a gradient foam structure would elicit a favorable response to chondrocytes or osteoblasts depending on the spatial location.

[0033] Further, the purpose of a functional gracient is to more evenly distribute the stresses across a region through which mechanical and/or physical properties are varying and thereby alleviate the stress concentrating effects of a sudden interface. This more closely resembles the actual biological tissues and structures, where structural transitions between differing tissues such as cartillage and bone are gradual. Therefore, it is an object of the present invention to provide an implant with a functional gradient between malerial phases. The present invention provides a multi-phasic functionally graded bicabsorbable implant with attachment means for use in surgical repair of estecohordral defects or sites of osteoarthrifis. Several patents have proposed systems for repairing cartilage that could be used with the present inventive pricus scafficids. For example, U.S. Patent 5,769,899 describes a device for repairing cartilage defects and U.S. Patent 5,713,974 describes securing cartilage repair devices with bone anchors (both hereby incorporated herein by reference).

Bone

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[0034] Gradient structures naturally occur for the bone / cartilage interface. In a study in our laboratories, we have demonstrated that material differences significantly influence cell function. In initial and long-term response of primary

cateoblasts to polymer (ilins (95.5 L-lacide-co-glycolide copolymer, 90/10 glycolide-co-(L)lacide copolymer, 95/5 L-lacide-co-explorations copolymer, 75/26 glycolide-co-(L)lacidic copolymer and 40/60 reagnolactone-co-(L)lacidic copolymer and 40/60 reagnolactone-co-(L)lacidic copolymer and kinited meshes (95/5 (L)lacide co-glycolide and 90/10 glycolide-co-(L)lacidic ecopolymers) were evaluated in vitro. The results demonstrated that osteoblasts attached and proliferated wellon all the biodegradable polymer films and meshes following 6-day incubation. None of the tested polymer films, except a 40/60 c-coprolactone-co-(L) lacidic copolymer film, demonstrated significant enhancement in differentiation of primary rat osteoblasts as compared to issue culture polystyrens (control). Films made of 40/60 c-sprolactone-co-(L) lacidic promoted enhanced differentiation of cultured ceteoblasts as demonstrated by increased alkaline phosphatase activity and osteoclacin mRNA expression as compared to the other films and TCPS. Hence, it is clear that different absorbable materials will significantly alter cell function and differentiation. By identifying the optimal materials for cell growth and differentiation a composite materials with a gradient composition could be utilized to optimize tissue regeneration with different cell types in the same scafflot.

[0055] Therefore, for bone repair or regeneration devices or scalloidings, a device made from a homopolymer, co-polymer (random, block, segmented block, tappered blocks, graft, triblock, etc.) having a linear, branched or star structure containing a caprolactone is especially preferred. Currently preferred are alphatic polyester copolymers containing in the range of from about 30 weight percent to about 99 weight percent e -caprolactone. Suitable repeating units that may be copolymerized with e-caprolactone reparried to a ref well known in the art. Suitable commonres that may be copolymerized with e-caprolactone include, but are not limited to lactic acid, lactide (including L., D. meso and D.L mixtures), glycolic acid, glycolide, p-dioxanone (1,4-dioxan-2-one), trimethylene carbonate (1,3-dioxan-2-one), &-valerolactone, @-butylene carbonate, etc., dioxan-2-one, including lactione, acid ethylopropiolactone, ethylene carbonate, ethylene carbonate, glycolic acid, glycolic pole, glyco

[0036] Preferred medical devices or tissue scaffoldings for bone tissue repair and/or regeneration containing bloebsorbable polymers made from e-captrolaction include but are not limited to the porcus beam scaffoldings (such as described in this application), Biorous three dimensional, spun, nonwoven, woven, knitted, or braided tissue scaffoldings, composite or cinciniation centriforcing bibers, matrices and combinations thereof.

Skin

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[0037] Another example of a tissue that has a gradient structure is skin. The basic structure of skin has two distinct. but well integrated layers where the thickness of each layer varies at different locations of the body. The outer layer or epidermis, is avascular and mainly consists of keratinocytes with smaller numbers of immune cells (Langerhan cells) and plamented cells (melanocytes). The keratinocytes produce keratin fibers and comeocyte envelopes, which gives the epidermis its durability and protective capabilities. The development of these structures is completely dependent upon the differentiation state of the epidermis. The epidermis forms a stratified epithelium, with different protein expression patterns, as the cells move further away from the basement membrane. This stratified layer of differentially expressing cells must be formed for maintenance of epidermal function. Below the epidermis is the dermis, which is a dense irregular connective tissue that is highly vascular. This layer is heavily populated with collageneic and elastic fibers, which give it its exceptional elasticity and strength. Fibroblasts are the main cell types in this layer. Between these two layers is the basement membrane, which serves as the site of attachment for epidermal cells and serves also to regulate their function and differentiation. The layer of keratinocytes, which attaches directly to the basement membrane, are cuboidal in shape and highly aligned. This attachment and architecture are critical requirements driving the ultimate production of the higher squamous structures in the epidermis. The basal layer provides a source of precursor cells for repair and replacement of the epidermis. The squamous layers provide strength and resistance to insult and infection.

[D038] Any material used for replacement of skin must be able to entice invasion of throblasts or other cells necessary to produce the demail components of the healed lissue. Additionally, the material must not inhiblt, and preferably should enhance, the rate of re-epithelialization in such a tashion that a discreet, epidermal basal layer is formed. Materials that permit invasion into the scaffold by migrating keratinocytes can produce partially differentiated cells. Consequently, control of access of particular cell types and a porous design that facilitates the regeneration of the natural tissue can have functional benefits. Now refer to Figures 9A, 9B and 9C which illustrates the microstructure of this foam scaffold figures 10 (100X magnification) and 11 (40X magnification composite provide proture) provide photomicrographic evidence of the invasion of libroblasts, macrophages, macrophage signal colls and ondothalistike cells into the a 0.5 mm foam. The foam tissue scaffolding 101 shown in both pictures was a \$0.50 blend of c-caprolactone-co-glycolide copolymer and c-caprolactone-co-alcide copolymer (made as described in Example 7). The pictures were taken at 8 days after implantation in 1.5 cm X 1.5 cm X 0.2 cm excisional wound model in a Yorkshire pig model. Complete incorporation of the matrix into the granulation tissue bed is evident in both pictures. The dense fibrous tissue above the foam tissue scaffolding appears to provide a suitable substate for the over growth of epidermis. PDGF was incorporated into the

foam tissue scaffolding shown in Figure 11. In compromised wound healing models the addition of a growth factor such as PDGF may in fact be necessary.

[0039] From our initial studies it appears that it is desirable to use as a skin scaffold a loam tissue scaffold having a thickness of from about 150 jum to about 3 mm, preferably the thickness of the foam may be in the range of from about 900 jum to loabout 1500 jum and most preferably about 500 to about 1000 jum. Clearly different skin injuries (i.e. diabetic ulces, venous stasis ulcers, decubitis ulcers, bums etc.) may require different loam thickness. Additionally, the patient condition may necessitate the incorporation of growth lactors, antibiotics and antifungal compounds to facilitate wound healing.

Vascular grafts

[0040] The creation of tubular structures with gradients may also be of interest. In vascular grafts, having a tube with pores in the outer diameter which transitions to smaller pores on the inner surface or visa versa may be useful in the culturing of endothetial cells and smooth muscle cells for the tissue culturing of vessels.

[0041] Multilayered tubular structures allow the regeneration of lissue that mimics the mechanical and/or biological characteristics of bloodyesels will have utility as a vascular grafts, Concentric layers, made from different compositions under different processing conditions could have taisoed mechanical properties, buebasoption properties, and tissue ingrowth rates. The niner most, or turninal layer would be optimized for endothelialization through control of the process of the surface and the possible addition of a surface treatment. The outermost, or adventitual layer of the vascular graft would be tailored to induce tissue ingrowth, again by optimizing the procesity (percent procesty, pore size, pore stage shape and pore size distribution) and by incorporating bloacelive factors, pharmaceutical agents, or cells. There may or may not be a barrier layer with low porosity between these two procus layers to increase strength and decrease leaders.

Composition of foams

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[0042] A variety of absorbable polymers can be used to make foams. Examples of suitable biocompatible, bioabsorbable polymers that could be used include polymers selected from the group consisting of aliphatic polyesters, poly (amino acids), copoly(ether-esters), polyalkytenes oxalates, polyamides, poly(iminocarbonates), polyorthoesters, polyoxaesters, polyamidoesters, polyoxaesters containing amine groups, poly(anhydrides), polyphosphazenes, biomolecules and blends thereof. For the purpose of this invention aliphatic polyesters include but are not limited to homopolymers and copolymers of lactide (which includes lactic acid, D-, L- and meso lactide), glycolide (including glycolic acid), ε-caprolactone, p-dioxanone (1.4-dioxan-2-one), trimethylene carbonate (1,3-dioxan-2-one), alkyl derivatives of trimethylene carbonate, δ-valerolactone, β-butyrolactone, γ-butyrolactone, ε-decalactone, hydroxybutyrate (repeating units), hydroxyvalerate (repeating units), 1,4-dioxepan-2-one (including its dimer 1,5,8,12-letraoxacyclotetradecane-7,14-dione), 1,5-dioxepan-2-one, 6,6-dimethyl-1,4-dioxan-2-one 2,5-diketomorpholine, pivalolactone, alpha, alpha-diethylpropiolactone, ethylene carbonate, ethylene oxalate, 3-methyl-1,4-dioxane-2,5-dione, 3,3-diethyl-1,4-dioxan-2.5-dione, 6,8-dioxabicycloctane-7-one and polymer blends thereof. Poly(iminocarbonate) for the purpose of this invention include as described by Kemnitzer and Kohn, in the Handbook of Biodegradable Polymers, edited by Domb, Kost and Wisemen, Hardwood Academic Press, 1997, pages 251-272. Copoly(ether-esters) for the purpose of this invention include those copolyester-ethers described in "Journal of Biomaterials Research", Vol. 22, pages 993-1009, 1988 by Cohn and Younes and Cohn, Polymer Preprints (ACS Division of Polymer Chemistry) Vol. 30(1), page 498, 1989 (e.g. PEC/PLA). Polyalkylene oxalates for the purpose of this invention include Patent Nos. 4,208,511; 4,141,087; 4,130,639, 4,140,678, 4,105,034; and 4,205,399 (incorporated by reference herein). Polyphosphazenes, co., ter- and higher order mixed monomer based polymers made from L-lactide, D,L-lactide, lactic acid, glycolide, glycolic acid, para-dioxanone, trimethylene carbonate and s-caprolactone such as are described by Allcock in The Encyclopedia of Polymer Science, Vol. 13, pages 31-41, Wiley Intersciences, John Wiley & Sons, 1988 and by Vandorpe, Schacht, Dejardin and Lemmouchi in the Handbook of Biodegradable Polymers, edited by Domb, Kosl and Wisemen, Hardwood Academic Press, 1997, pages 161-182 (which are hereby incorporated by reference herein). Polyanhydrides from diacids of the form HOOC-C₆H₄-O-(CH₂)_m-O-C₆H₄-COOH where m is an integer in the range of from 2 to 8 and copolymers thereof with aliphatic alpha-omega diacids of up to 12 carbons. Polyoxaesters, polyoxaamides and polyoxa aesters containing amines and/or amido groups are described in one or more of the following U.S. Patent Nos. 5,464,929; 5,595,751; 5,597,579; 5,607,687; 5,618,552; 5,620,698; 5,645,850; 5,648,088; 5,698,213; 5,700,583; and 5,859,150 (which are incorporated herein by reference). Polyorthoosters such as those described by Heller in Handbook of Biodegradable Polymers, edited by Domb, Kost and Wisemen, Hardwood Academic Press, 1997, pages 99-118 (hereby incorporated herein by reference).

[0043] Currently aliphatic polyesters are the absorbable polymers that are preferred for making gradient foams. Alphatic polyesters can be homopolymers, copolymers (random, block, segmented, tappered blocks, graft, triblock, etc.) having a linear, branched or star structure. Preferred are linear copolymers. Suitable monomers for making aliphatic homopolymers and copolymers may be selected from the group consisting of, but are not limited, to lactic acid, lactide (including L., D., meso and D.L mixtures), glycolic acid, glycolide, r-caprolactione, p-dioxanone (1,4-dioxano-2-one), trimethylene carbonate (1,3-dioxano-2-one), delta-valerolactione, beta-butyrolactione, epsion-decalactione, 2,5-diletomorpholine, phyalolactione, alpha, alpha-diethylproprolactione, ethylene carbonate, ethylene oxalate, 3-me-thyl-1,4-dioxano-2,5-dione, 3,3-diethyl-1,4-dioxano-2,5-dione, 3,3-diethyl-

[0044] Elastomeric copolymers also are particularly useful in the present invention. Suitable bioabsorbable biocompatible elastomers include but are not limited to those selected from the group consisting of elastomeric copolymers of ε-caprolactone and glycolide (preferably having a mole ratio of ε-caprolactone to glycolide of from about 35:65 to about 65:35, more preferably from 45:55 to 35:65) elastomeric copolymers of s-caprolactions and lactide, including Llactide, D-lactide blends thereol or lactic acid copolymers (preferably having a mole ratio of s-caprolactions to lactide of from about 35:65 to about 65:35 and more preferably from 45:55 to 30:70 or from about 95:5 to about 95:15) elastomeric copolymers of p-dioxanone (1,4-dioxan-2-one) and lactide including L-lactide, D-lactide and lactic acid (preferably having a mole ratio of p-dioxanone to lactide of from about 40:60 to about 60:40) elastomeric copolymers of Ecaprolactone and p-dioxanone (preferably having a mole ratio of ε-caprolactone to p-dioxanone of from about from 30: 70 to about 70:30) elastomeric copolymers of p-dioxanone and trimethylene carbonate (preferably having a mole ratio of p-dioxanone to trimethylene carbonate of from about 30:70 to about 70:30), elastomeric copolymers of trimethylene carbonate and glycolide (preferably having a mole ratio of trimethylene carbonate to glycolide of from about 30;70 to about 70:30), elastomeric copolymer of trimethylene carbonate and lactide including L-lactide, D-lactide, blends there of or lactic acid copolymers (preferably having a mole ratio of trimethylene carbonate to lactide of from about 30:70 to about 70:30) and blends thereof. Examples of suitable bioabsorbable elastomers are described in U.S. Patent Nos. 4,045,418; 4,057,537 and 5,468,253 all hereby incorporated by reference. These elastomeric polymers will have an inherent viscosity of from about 1.2 dL/g to about 4 dL/g, preferably an inherent viscosity of from about 1.2 dL/g to about 2 dL/g and most preferably an inherent viscosity of from about 1.4 dL/g to about 2 dL/g as determined at 25°C in a 0.1 gram per deciliter (g/dL) solution of polymer in hexafluoroisopropanol (HFIP).

[O045] Proferably, the elastomers will exhibit a high percent elongation and a low modulus, while possessing good tensile strength and good recovery characteristics. In the preferred embodiments of this invention, the elastomer from which the loams are formed will exhibit a percent elongation greater than about 200 percent and preferably greater than about 500 percent. There properties, which measure the degree of elasticity of the bioabsorbable elastomer, are achieved while maintaining a tensile strength greater than about 500 ps, preferably greater than about 500 bs/mch, preferably greater than about 500 bs/mch.

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[0046] The polymer or copolymer suitable for forming a gradient foam for tissue regeneration depends on several factors. The chemical composition, spatial distribution of the constituents, the molecular weight of the polymer and the degree of crystallinify all dictate to some extent the *in-vitro* and *in-vivo* behavior of the polymer. However, the selection of the polymers to make gradient foams for tissue regeneration largely depends on (but not limited to) the following factors: (a) bioabsorption (or bio-degradation) kinetics; (b) in-vivo mechanical performance; and (c) cell response to the material in terms of cell attachment, proliteration, migration and differentiation and (d) biocompatibility.

[0047] The ability of the material substrate to resorb in a timply tashion in the body environment is critical. But the differences in the absorption time under in vivioconditions can also be the basis for combining two different copolymers. For example a copolymer of 35.65 e-caprolactone and glycolide (a relatively fast absorbing polymer) is blended with 40:50 e-caprolactone and (L)lactide copolymer (a relatively slow absorbing polymer) to form a foam. Such a fearn could have several different physical structures depending upon the processing lechnique used. The two constituents can be either randomly inter-connected bicontinuous phases, or the constituents can have a gradient through the thickness or a laminate type composite with a well integrated interface between the two constituent layers. The microstructure of these foams can be optimized to regenerate or repair the desired anatomical features of the lissue that is being engineered.

[0048] One preferredembodiment of the present invention is to use polymer blends to form structures which transition from one composition to involve composition in a gradient like architecture. Forem having this gradient architecture are particularly advantageous in tissue engineering applications to repair or regenerate the structure of naturally co-curring tissues such as cartilage (articular, meniscal, septal, tracheal etc.), esophiaguses, skin, bone and vascular tissue. For example by blending an elastiomer of e-caprolactione-co-plycolide with e-caprolactione-co-lactide (i.e. with a mole rails of about 5.95) a foam may be formed that transitions from a softer spongy foam to a stifler more rigid foam similar to the transition from cartilage to bone. Clearly other polymer blends may be used for similar gradient offsicts or to provide different gradients such as different absorption profiles, stress response profiles, or different degrees of elasticity. Additionally, these foams can be used for organ repair replacement or regeneration strategies that may benefit from these unique scaffolds, including but are not limited to, spine disc, dura, nerve tissue, liver, pancreas, kidney, bladder, tendone, ligaments and breast tissues.

[0049] These elastomeric polymers may be foamed by lyophilization, supercritical solvent foaming (i.e., as described

in EP 464,163 B1), gas injection extrusion, gas injection molding or casting with an extractable material (i.e., selts, sugar or any other means known to those skilled in the ani). Currently it is preferred to prepare bioabsorbable, biocompatible elastometric foams by tyophilization. Suitable methods for tyophilizing elastometric polymers to form foams is described in the Examples and in the copending patent application entitled. "Process for Manufacturing Biomedications of B

[0050] The learns that are made in this invention are made by a polymer-solvent phase separation technique with modifications to the prior art that unexpectedly creates gradients in the loam structure. Generally, a polymer solution can be separated into two phases by any one of the four techniques. (a) thermally induced gelation/crystalization, (b) non-solvent induced separation of solvent and polymer phases; (c) chemically induced phase separation, and (d) thermally induced spinodal decomposition. The polymer solution is separated in a controlled menner into either two distinct phases or two bicontinuous phases. Subsequent removal of the solvent phase usually leaves a porous structure of density less than the bulk polymer and pores in the micrometer ranges (ref. "Microcellular foams viaphase separation" by A. T. Young, J. Vac. Sci. Technolol. A 4(3), May/Jun 1986). The steps involved in the preparation of these foams consists of choosing the right solvents for the polymers that needs to be hyphilized and preparing a homogeneous solution. Next, the polymer solution and vacuum drying step removes the solvent by sublimation and/or drying leaving a porous polymer structure or an interconnected open coll prorous fear.

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[0051] Suitable solvents that should be generally suited as a starting place for selecting a solvent for the preferred absorbable alliphatic polyesters include but are not limited to solvents selected from a group consisting of formic acid, ethyl formate, acetic acid, hoxafturorisepropanol (HFIP)-cyclic ethers (e. THE, DMF, and PDO), acetione, acetales of C2 to C5 alcohol (such as ethyl acetate and t-butylacetate), glyme (i.e. monoglyme, ethyl glyme, diglyme, ethyl glyme, diglyme, activated and the service of t

[0052] Accordingly, as will be appreciated, the applicable polymer concentration or amount of solvent, which may be utilized, will vary with each system. Suitable phase diagram curves for several systems have already been developed. However, if an appropriate curve is not available, this can be readily developed by known techniques. For example, a suitable technique is self orth in Smolders, van Aartsen and Steenbergen, Kolloid-Z. u. Z. Polymere, 243, 14 (1971), as a general guideline the amount of polymer in the solution can vary from about 0.5% to about 90% and preferably will vary from about 0.5% to about 90% and preferably preferably preferably preferably preferably preferably prefer

[0053] Additionally, solids may be added to the polymer-solvent system. The solids added to the polymer-solvent system preferably will not react with the polymer or the solvent. Suitable solids include materials that promote tissue regeneration or regrowth, buffers, reinforcing materials or porosity modifiers. Suitable solids include, but are not limited to, particles of demineralized bone, calcium phosphate particles, or calcium carbonate particles for bone repair, leachable solids for pore creation and particles of bioabsorbable polymers not soluble in the solvent system as reinforcing or to create pores as they are absorbed. Suitable leachable solids include but are not limited nontoxic leachable materials selected from the group consisting of salts (i.e. sodium chloride, potassium chloride, calcium chloride, sodium tartrate, sodium citrate, and the like) biocompatible mono and disaccharides (i.e. glucose, fructose, dextrose, mallose, lactose and sucrose), polysaccharides (i.e. starch, alginate), water soluble proteins (i.e. gelatin and agarose). Generally all of these materials will have an average diameter of less than about 1mm and preferably will have an average diameter of from about 50 to about 500 µm. The particles will generally constitute from about 1 to about 50 volume percent of the total volume of the particle and polymer-solvent mixture (wherein the total volume percent equals 100 volume percent). The leachable materials can be removed by immersing the foam with the leachable material in a solvent in which the particle is soluble for a sufficient amount of time to allow leaching of substantially all of the particles, but which does not dissolve or detrimentally alter the foam. The preferred extraction solvent is water, most preferably distilled-delonized water. This process is described in U.S. patent No. 5,514,378 hereby incorporated herein by reference (see column 6). Preferably the foam will be dried after the leaching process is complete at low temperature and/ or vacuum to minimize hydrolysis of the foam unless accelerated absorption of the foam is desired.

[0054] After the polymer solvent mixture is formed the mixture is then solidified. For a specific polymer-solvent system, the solidification point, the melt temperature and the apparent glass transition of the polymer-solvent system can be determined using standard differential scanning calorimetric (DSC) techniques. In theory, but in no way limiting the scope of the present invention, it is believed that as a polymer solvent system is cooled down an initial solidification occurs at about or below the freezing point of the solvent. This corresponds to the freezing of a substantial portion of

the solvent in the system. The initial freezing appears as a first exothermic peak. A second freezing point occurs when the remaining solvent associated with the polymer solidifies. The second freezing point is marked by a second exothermic peak. The apparent Tg is the temperature at which the fully frozen system displays the first endothermic shift on reheating.

10055] An important parameter to control is the rate of freezing of the polymer-solvent system. The type of pore morphology that gets locked in during the freezing step is a function of the solution thermodynamics, freezing rate, temperature to which it is cooled, concentration of the solution, homogeneous or heterogeneous nucleation etc. Detailed description of these phase separation phenomenon can be found in the references provided herein ("Microcellular foams via phase separation" by A. T. Young, J. Vac. Sci. Technol. 4 4(3), MayJun 1986; and Thermodynamics of Formation of Porous Poymeris Membrane from Solutions* by S. Matsuda, Polymer J. Vol. 23, No. 5, pp 435-444, 1991). [2056]

The polymer solution previously described can be solidilitied in a variety of manners such as placing or injecting the solution in a mold and cooling the mold in an appropriate bath or on a retrigerated shelf. Alternatively, the polymer solution can be atlomized by an atomizer and sprayed onto a cold surface causing solidification of the spray layer by layer. The cold surface can be a medical device or part thereof or a film. The shape of the solidifical spray will be shape of the solution spray shapes, because and other processes the teams can be made or molded in a variety of shapes and sizes (i. e. tubular shapes, branched tubular shapes, spherical shapes, hemispherical shapes, in furst conical shapes, pyramidal shapes, elipsoidal shapes (i.e. kidney shaped, to combination the recol.)

[0057] Alternatively, another method to make shaped (owned parts is to use a cold finger (a metal part whose surface represents the inside of the part we want to fabricate). The cold finger is dipped into a matture of polymer in an appropriate solvent and removed. This is much like dipping an ice cream pop into warm chocolate that freezes to a hard, cold skin, or dipping a form into a latex of rubber to form gloves or condoms. The thickness and morphology of the foam produced are a function of the temperature, dwell time and whitidrawal rate of the cold finger in the mixture. Longer dwolt, coldor finger and slower withdrawal will produce a thicker coating. After withdrawal, the cold finger is placed on a fixture of large themal mass that is in contact with the refrigerated tray of the hypohilizer. From this point the primary and secondary drying processes are as described above. This method is particularly well suited to making these, branched tubular structures or sleeves that may be shaped to fit devices or portions of an animal's anatomy (for repair, regeneration or augmentation of tissue).

20 [ODS8] Additionally, the polymer solution can be solidified with various inserts incorporated with the solution such as films, scrims, weven, nonwoven, knitted or braided textile structures. Additionally, the solution can be prepared in association with another structure such an orthopedic implant (e.g. screws, pins, nails, and plates) or vascular or branched tubular construct (as a scatilotif or a vascularized or ducted organ). These inserts will be made of at least one biocompatible material and may be non-absorbable, absorbable or a combination thereof.

[0059] The polymer solution in a mold undergoes directional cooling through the wall of the mold that is in contact with the freeze driver shelf, which is subjected to a thermal cycle. The mold and its surface can be made from virtually any material that does not interfere with the polymer-solvant system, though it is preferred to have a highly conducting material. The heat transfer front moves upwards from the lyophilizer shelf through the mold wall into the polymer solution. The instant the temperature of the mixture goes below the gellation and/or freezing point the mixture also phase separates.

[0060] The morphology of this phase separated system is locked in place during the freezing step of the lyophilization process and the creation of the open pores is initiated by the onset of vacuum drying resulting in the sublimation of the solvent. However, the mixture in container or mold that is cooled from a heat sink will solidity injor to completely freezing. Although the mixture may appear solid, initially there appears to be some residual solvent associated with the polymer that has not cystallized. It is theorized, but in no way limiting the present invention, that a freezing from moves through the mixture from the heat sink to complete the solidification after the mixture has apparently solidified. The material in front of the freezing front at a given time will not be as cold as the material behind the front and will not be in a completely fozzen state.

[0061] We have discovered that if a vacuum is applied to the apparently solid polymer-solvent mixture immediately after it appears to solidify, a foam with a gradient structure having variable pore size and structure as well as channels can be created. Therefore, timing of the creat of the sublimation process (by pressure reduction i.e. vacuum dying) is a critical step in the process to create transitions in the structure. The timing of the onset of sublimation will be affected by the hickness of the foam boingmade, concentration of the solution, rate of heat transfor, and directionation of the solution, rate of heat transfor, and directionation of the solution, rate of heat transfor and characterized for specific polymer-solvent systems by using thermocouples and monitoring the heat transfer rates of the fearms at various depths and locations in the device being foamed. By controlling the sublimation process, structures with a gradient in pore morphology and anisotropy may be created. This process can lead to the creation of microstructures that mimic tissues such a scrittinge, bone and skin. For example, channels will generally be formed if a vacuum is putted mixed.

diately after the solution apparently solidities. However, if the same solution is allowed to solidify further the foam will have larger pores closer to the surface where the vacuum is being drawn (opposite the heat sink) and smaller pores closer to the heat sink.

[0062] This process is the antitheses of the prior art process that focused on creating foams with a uniform microstructure (randomy interconnected pores), whereby the whole solution is completely frozen. And vascum drying is applied only after a considerable amount of time is given for the completion of desired phase separation (U.S. Patients, 5,765,792 (Brakke), 5;133,755 (Brakke), 5;376,413 (Walter, et al.), 5,607,474 (Athanasiou, et al.); 5,608,091 (Leong, et al.), 5,507,355 (Shalbey, et al.); and European disclosures E0724989 (Hinsch) and EPA 5941449 (Totalvara)

[0063] Foams with various structures are shown in Figures 2, 3, and 4. For example, as shown in Figure 3 the orientation of the major axis of the porces may be changed from being in the same plane as the foam to being oriented perpendicular to the plane of the foam. By way of theory, but in no way limiting the ecope of this invention, it is betied that this in conventional foam processing as the solvent crystallizes at reczing front moves through the solution solid-living the solution in crystalline layers parallel to the freezing front. However, if a vacuum is publicable force the solution completely freezes, the morphology of the foam results in pores being formed generally aligned parallel to the vacuum source. As is illustrated in Figure 3.

[0064] As can be seen from Figure 3 the pore size can be varied from a small pore size generally between about 10 µm and about 60 µm to a larger size of from about 60 µm to about 200 µm in a porous gradient loam. Again this results from pulling a vacuum on the apparently solidified solution before its completely solidified. The polymer concentration in the solution and the cooling rates are also important parameters in controlling the cell size. Ideally the foam structure could be created to serve as a template to restore human tissue junctions such as the cartilage to bone junction present in joints. This foam would progress form a small round pores to larger column-like (i.e. with a diameter to length ratio of at least 2 to 1 1) poes. Additionally, the stiffness of the foam can controlled by the foams structure or blending two different polymers with different Young's modult.

[0065] Foams can also have channels as is illustrated in Figure 2. The channels formed by this process may traverse the thickness of the loam and generally range in diameter from about 30 to about 200 µm in diameter. The channels generally are at least two times the channels average diameter and preferably are at least four times the channels average diameter and most preferably at least eight times the channels average diameter from the channels average diameter of course will be selected based on the desired functionality of the channel such as cell invasion, nutrient diffusion or as a vernue for vascularization.

[0065] One skilled in the art can easily visualize that such a directionality can be created in any three dimensions by designing an appropriate motel and subjecting the walls of such a mold to different temperatures if needed. The following types of gradent structures can be made by variation in the pore size and/or shape through the thickness with a uniform composition; pores of one type and size for a certain thickness followed by another type and size of pores, etc. composition; pores of the size of an extension from positional gradent with predominantly one composition or one side and another one one the other with a transition from one entilly to the other, a thick skin comprising low poresity of low pore size layer followed by a large pore size region; cleans with vertical pores with a spatial organization these vertical pores can act as channels for nutrient diffusion the making of these in 3D molds to create 3D fearns with the desired microstructure combinations of compositional architectural gradient. Generally the fearns formed in containers or molds will have a thickness in the range of from about 0.25mm to about 50mm. This will have a thickness of from about 0.5mm to about 50mm. Though containers or molds with the residual solvent content may be quite long, the final fearn structures may be more difficult to control and the residual solvent content may be higher.

[0067] As previously described the inventive process cycle for producing biocompatible foam is significantly reduced by performing the sublimation step above the apparent glass transition temperature and below the solidification temperature.) The combined cycle time of treating performing the sublimation step above the apparent glass transition temperature or the muture (preferably just below the solidification temperature). The combined cycle time of treating primary drying + secondary drying) is much faster than is described in the prior art. For example, the combined cycle to aliphatic polyseters using vostatie solvents is generally less than 72 hours, preferably less than 48 hours, more preferably less than 24 hours and most preferably less than 10 hours. In fact the combined cycle can be performed with some aliphatic polyseters in less than 3 har jor foams of thickness farm or less, less than 6 hrs for foams of thickness around 2 mm. Compare this with prior at which is typically 72 hours or greater. The residual solvent concentrations in these foams made by this process will be very fow As described for aliphatic polyseters foams made using 1.4-dioxane was less than 10 ppm (parts per million) more preferably less than 1 ppm and most preferably less than 100 ppb (parts per billion).

[0068]. One skilled in the art can easily visualize that such a directionality can be created in any three-dimensions by designing an appropriate mold and subjecting the walls of such a mold to different temperatures if needed. The following types of gradient structures can be made by this invention.

variation in the pore size and/or shape through the thickness with a uniform composition.

- 2. pores of one type and size for a certain thickness followed by another type and size of pores, etc.
- compositional gradient with predominantly one composition on one side and another composition on the other with a transition from one entity to the other
- 4. a thick skin comprising low porosity of low pore size layer followed by a large pore size region
- 5. foams with vertical pores with a spatial organization...these vertical pores can act as channels for nutrient dif-
- the making of these in three-dimensional molds to create three-dimensional foams with the desired microstructure.
 - 7, combinations of compositional and architectural gradient

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[0069] Additionally, various proteins (including short chain péptides), growth agents, chemotatic agents and therapeutic agents (antibiotics, analgesies, antilinflammations, anti-rejection (e.g., immunosuppressants) and anticancer drugs), or ceramic particles can be added to the foams during processing, advobed onto the surface or back filled into the foams after the loams are made. For example, the pores of the foam may be partially or completely filled with biocompatible resorbable synthetic polymers or biopolymers (such as collagen or elastin) or biocompatible ceramic materials (such as hydroxyapatite) and combinations thereof (that may or may not contain materials that promote tissue growth through the device). Suitable materials include but are not limited to autograft, allograft, or xenograft bone, bone marrow, morphogenic proteins (BMPS), epidermal growth factor (EGF), libroblast growth factor (FGF), platelet derived growth factor (FDGF), insulin derived growth factor (IGF-1 and IGF-1I), transforming growth factors (TGF-8), vascular endoblestil growth factor (VEGF) or other osteoinductive or osteoconductive materials known in the at 18 opolymers could also be used as conductive or chermotactic materials, or as delivery vehicles for growth factors. Examples could be recombinant or animal derived college or eleation or hyaluronic acid. Bioactive ceatings or surface treatments could also be attached to the surface of the materials. For example, bioactive peptide sequences (RGD's) could be attached to lacilitate protein adsorption and subsequent cell tissue attachment. Therapeutic agents may also be delivered with these foams.

[0079] In another embodiment of the present invention, the polymers and blends that are used to form the foam can contain therapeutic agents. To form these foams, the previously described polymer would be mixed with a therapeutic agent prior to forming the foam or loaded into the foam after it is formed. The variety of different therapeutic agents that can be used in conjunction with the foams of the present invention is vast. In general therapeutic agents which may be administered via the pharmaceutical compositions of the invention include, without timitation: antilitrative such as antibiotics and antiviral agents; chemotherapeutic agents (i.e. anticancer agents); anti-rejection agents; analyseis and analgesic combinations; anti-inflammatory agents, homones such as steroids; growth factor (bone morphogenic proteins (i.e. BMP 9-17), bone morphogenic-like proteins (i.e. GFD-5, GFD-7 and GFD-9), epidemial growth factor (EGFF), insufn fixe growth factor (GFF), insufn fixe growth factor (GFFF), insufn fixe growth factor (GFFFF), insufn f

[0071] Foams containing bio-active materials may be formulated by mixing one or more therapeutic agents with the polymer used to make the foam or with the solvent or with the solventor with the solventor with the solventor with the solventor with the polymer-solvent mixture and foamed. Alternatively, a therapeutic agent could be coated on to the foam preferably with a pharmaceutically acceptable carrier.

[0072] Any pharmaceutical carrier can be used that does not dissolve the foam. The therapeutic agents, may be present as a liquid, a finely divided solid, or any other appropriate physical form. Typically, but optionally, the matrix will include one or more additives, such as ditients, carriers, excipents, stabilizers or the like.

[0073] The amount of therapeutic agent will depend on the particular drug being employed and medical condition being trailed. Typleally, the amount of drug represents about 0.001 percent to about 70 percent, more typically about 0.001 percent to about 20 percent by weight of the matrix. The quantity and type of polymer incorporated into the drug delivery matrix will vary depending on the release profile desired and the amount of drug employed.

[0074] Upon contact with body fluids the drug will be released. If the drug is incorporated into the foam then as the foam undergoes gradual degradation (mainly through hydrolysis) the drug will be released. This can result in prolonged delivery (over, say 1 to 5,000 hours, preferable 2 to 800 hours) of effective amounts (say, 0,000 in mykyghour) to 10 mg/kg/hour) of the drug. This dosage form can be administered as is necessary depending on the subject being treated, the severity of the affliction, the judgment of the prescribing physician, and the like. Following this or similar procedures, those skilled in the art will be able to prepare a variety of formulations.

[0075] The foam may also serve as a scallfold for the engineering of tissue. The porous gradient structure would be conductive to growth of cells. As outlined in previous patents (Vacanti, U.S. 5,770,417), cells can be harvested from a patient (before or during surgery to repair the tissue) and the cells can be processed under startle conditions to provide

a specific cell type (i.e., pluripotent cells, stem cells or precursor cells such as the mesenchymal stem cells described in Capian, U.S. 5. 486,359, etc.). Suitable cell that may be contacted or seeded into the foam scaliolds include but are not limited to myocytes, adipocytes, libornyroblasts, ectodermal cell, muscle cells, cetodblast (i.e. bone cells), chondrocyte (i.e. cartilage cells), endothelial cells, fibroblast, pancreatic cells, hepatocyte, bile duct cells. bone marrow cells, neural cells, genitourinary cells (including nephritic cells) and combinations thereol. Various cellular strategies cells results and the cells stead with these scalfolds (i.e. autogenous, allogenic, xenogeneic cells etc.). The cells could also contain inserted DNA enceding a protein that could stimulate the attachment, proliferation or differentiation of tissue. The foam would be placed in cell culture and the cells seeded onto or into the structure. The foam would be maintained in a sterile environment and then implanted into the donor patient once the cells have invaded the microstructure of the device. The in vitro seeding of cells could provide for a more rapid development and differentiation process for the tissue. It is clear that cellular differentiation process for the tissue. It is clear that cellular differentiation process for the dissue.

[0076] The option for seeding different cell types into the different pore structures would be available to investigators. Schaufer et al., have demonstrated that different cell types (i.e. stromal cells and chondrocytes) can be cultured on different structures. The structures can be combined after a short period of time and the entire structure can be particulated back in cell culture so a biphasic issues structure can be generated for implantation. A gradient structure would also allow for co-cultured issue scatclos to be generated. (Schaefer, D. et al.) Additionally, radio-cpaque markers may be added to the foams to allow imaging after implantation. After a delined period of in vitro development (for example weeks), the tissue empineered implant would be harvested and implanted into the patient. If an acellular strategy is pursued, then the sterile acellular scanfolds would be used to replace damaged or traumatized itssue.

[0077] The foam scaffolds of the present invention may be sterilized using conventional sterilization process such as radiation based sterilization (i.e. gamma-ray), lemical based sterilization (ethylene oxide) or other appropriate procedures. Preferably the sterilization process will be with ethylene oxide at a temperature between 52-55°C for a time of 8 hours or less. After sterilization the foam scaffolds may be packaged in an appropriate sterilization transfer and use in hospitals and other health care facilities.

[0078] The following examples are illustrative of the principles and practice of this invention, although not limited thereto. Numerous additional embodiments within the scope and spirit of the invention will become apparent to those skilled in the art.

Examples

[0079] In the examples which follow, the polymers and monomers were characterized for chemical composition and purity (NMR, F.T-IR), thermal analysis (DSC), molecular weight (inherent viscosity), and baseline and in <u>vitro</u> mechanical properties (Instron stress/strain).

[0080] ¹H MMR was performed on a 300 MHz MMR using CDCl₃ or HFAD (hoxafluoroacetone sesqua deutrium oxide) as a sobrent. Themal analysis of segmented polymers and monomers was performed on a Dupont 912 Differential Scanning Calorimater (ISC). Inherent viscosities (I.V, d./g) of the polymers and cooptupers were measured using a 50 bore Cannon-Ubbelhock dilution viscometer immersed in a thermostatically controlled water bath at 25°C utilizing chloroform or hexafluorosopropanol (HFIP) as the solvent at a concentration of 0.1 gdt.

[0081] In these examples certain abbreviations are used such as PCL to indicate polymerized ε-caprolactone, PGA to indicate polymerized glycolide, PLA to indicate polymerized (L)lactide. Additionally, the percentages in Iront of the copolymer indicates the respective mole percentages of each constituent.

Example 1

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Preparation of a foam with random microstructure (no preferred architecture)

Step A. Preparing 5% wt./wt. homogeneous solution of 35/65 PCL/PGA in 1,4-Dioxane

[0082] A 5 % xd./wl. polymer solution is prepared by dissolving 1 part of 35/65 PCLPGA with 19 parts of the solvent - 1,4-dioxane. The 35/65 PCLPGA copolymer was made substantially as described in Example 8. The solution is prepared in a Itask with a magnetic stir bar. For the copolymer to dissolve completely, it is recommended that the mixture is gently heated to 60 ± 5°C and continuously stirred for a minimum of 4 hours but not exceeding 8 hours. A clear homogeneous solution is then obtained by filtering the solution through an extra coarse porcesty litter (Pyrex brand extraction thimble with Initied disc) using dry nitrogen to help in the filtration of this viscous solution. Step B. Lyophilization

[0083] A laborationy scale hyophilizer. Freezemobile 6 of VIRTIS was used in this experiment. The Irreeze dryer is powered up and the shell chamber is maintained at 20°C under dry nitrogen for approximately 30 minutes. Thermocouples to monitor the shell temperature are attached for monitoring. Carelluly fill the homogeneous polymer solution prepared in Step A. into the molds just before the actual start of the cycle. A glass mold was used in this example but a mold made of any material that is inert to 1.4 cilcoxine; has good heat transfer characteristics; and has a surface that enables the easy removal of the Ioam can be used. The glass mold or dish used in this example weighed 620 grams, was optical glass 5 nm hick, and cylindrical with a 21cm outer diameter and a 19.5cm inner diameter. The lip height of the dish was 2.5cm. Next the following steps are followed in a sequence to make a 2mm thick foam:

- (i). The glass dish with the solution is carefully placed (without tilting) on the shell of the lyophilizer, which is maintained at 20°C. The cycle is started and the shell temperature is held at 20°C for 30 minutes for thermal conditioning.
- (ii). The solution is then cooled to -5°C by cooling the shelf to -5°C.
- (iii) Alter 60 minutes of freezing at -5°C, a vacuum is applied to initiate primary drying of the dioxane by sublimation. One hour of primary drying under vacuum at -5°C is needed to remove most of the solvent. At the end of this drying stage typically the vacuum level reached about 50 mTorr or less.
- (iv). Next, secondary drying under a 50 mTorr vacuum or less was done in two stages to remove the adsorbed dioxane. In the lirist stage, the shelf temperature was raised to 5°C and held at that temperature for 1 hour. At the end of the first stage the second stage of drying was begun. In the second stage of drying, the shell temperature was raised to 20°C and held at that temperature for 1 hour.
- (v). At the end of the second stage, the lyophilizer is brought to room temperature and the vacuum is broken with nitrogen. The chamber is purged with dry nitrogen for approximately 30 minutes before opening the door

[0084] The steps described above are suitable for making foams that are about 2mm thick or less. As one skilled in the art would know, the conditions described herein are typical and operating ranges depend on several factors e.g.: concentration of the solution; cybmer molecular weights and compositions; volume of the solution; mold parameters; machine variables like cooling rate, heating rates; and the like. Figure 1 shows a SEM of a cross section of the foam produced following the process set forth in this example. Note the random microstructure (not a preferred architecture) of this loam.

Example 2

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Preparation of a foam with vertical channels

[0085] This example describes the making of a 35/65 PCL/PGA foam with vertical channels that would provide pathways for nutrient transport and guided tissue regeneration.

[0086] We used a FTS Dura Dry Freeze dryer with computer control and data monitoring system to make this foam. First step in the preparation of this foam was to generate a homogeneous solution. A 10% wt.Mt. homogeneous solution of 35/65 PCL/PGA was made in a manner similar to that described in Example 1, Step A. The polymer solution was carefully filled into a clish just before the actual start of the cycle. The dish weighed 620 grams, was optical glass 5.5mm thick, and cylindrical with a 21 cm outer diameter and a 19.5cm inner cliameter. The lip height of the dish was 2.5cm. Next the following steps are followed in sequence to make a Zmm thick foam with the desired architecture:

- (i). The solution filled dish was placed on the freeze dryer shelf that was precooled to -17°C. The cycle was started and the shelf temperature was held at -17°C for 15 minutes quenching the polymer solution.
- (ii). After 15 minutes of quenching to -17°C, a vacuum was applied to initiate primary drying of the dioxane by sublimation and hold at 100 milliTorr for one hour.
- (iii) Next, secondary drying was done at 5°C for one hour and at 20°C for one hour. At each temperature the vacuum level was maintained at 20 mTorr.
 - (iv). At the end of the second stage, the lyophitizer was brought to room temperature and the vacuum was broken

with nitrogen. The chamber was purged with dry nitrogen for approximately 30 minutes before opening the door.

[0087] Figure 2 is a SEM picture that shows a cross section of the foam with vertical channels. These channels run through the thickness of the foam.

Example 3

Architecturally gradient foam

[0088] This example describes the making of a toam that has a gradient in foam morphology as shown in Figure 3 using a 10% solution of 3865 ε-caprolactone-co-glycolide. The method used to make such a foam is similar to the description given in Example 2 with one difference. In step (ii) of the lyophilization process the time for which the solution is kept at the freezing step is 30 minutes.

[0089] Figure 3 is a scanning electron micrograph of a cross section of this foam. Note the variation in the pore size and pore shape through the thickness of the foam.

Example 4

Transcompositional foam

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[0090] This example describes the making of a foam that has a compositional gradient and not necessarily a morphological gradient. Such a foam is made from polymer solutions that have been made from physical mixtures of

25 Step A. Preparing a solution mixture of 35/65 PCL/PGA and 40/60 PCL/PLA in 1,4-Dioxane

[0061] In the preferred method the two separate solutions are first prepared (a) a 10% wtwt polymer solution of \$5/65 PCL/PGA and (b) a 10% wtwt 40/60 PCL/PLA. Once these solutions are prepared as described in Example 1, equal parts of each solution was poured into one mixing flast. The polymers used for make these solutions are described in Examples 8 and 3 Abmongeneous solution of this physical mixture was obtained by gently heating to 60 ± 5°C and continuously stirring using a magnetic sit has for approximately 2 hours.

Step B. Lyophilization cycle

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- 35 [0092] We used an FTS Dura Dry Freeze dryer with computer control and data monitoring system to make this foam. The first step in the preparation of such a loam was to generate a homogeneous solution as described in Step A. The solution was carefully filled into a dish just before the actual start of the cycle. The cylindrical glass dish weighed 117 grams, was optical glass 2.5mm thick and cylindrical with a 100mm outer diameter and a 95mm inner diameter. The lip height of the dish was 50mm. Next the following steps were followed in sequence to make a 25mm thick foam with the transcomosilional oradient:
 - (i). The solution filled dish was placed on the freeze dryer shelf and the solution conditioned at 20°C for 30 minutes. The cycle was started and the shelf temperature was set to -5°C with a programmed cooling rate of 0.5°C/min.
 - (ii). The solution was held at the freezing condition (-5°C) for 5 hours.
 - (iii). Vacuum was applied to initiate primary drying of the dioxane by sublimation and held at 100 milliTorrfor 5 hours.
 - (iv). Next, secondary drying was done at 5°C for 5 hours and at 20°C for 10 hours. At each temperature the vacuum level was maintained at 20 mTorr.
 - (v). At the end of the second stage, the lyophilizer was brought to room temperature and the vacuum was broken with nitrogen. The chamber was purged with dry nitrogen for approximately 30 minutes before opening the door.
- 5 [0093] The foam has a gradient in chemical composition which is evident from a close scrutiny of the foam wall morphobogy as shown in Figure 4, 5 and 6. The gradient in the chemical composition was further supported by NMR data as detailed below.
 - [0094] Foam sample produced by the above method and which was approximately 25 mm thick was characterized

for mole % composition. The foam sample is composed of a physical blend of PCL/PLA and PCL/PGA. Slices of the foam sample were prepared and analyzed to confirm that the material was a compositional gradient. The sample slices were identified as 1) foam 1A (top slice), 2) foam IB (top middle slice), 3) foam IC (bottom middle slice), 4) foam ID (bottom slice). The NMR sample preparation consisted of dissolving a 5mg of material into 300µL hexafluoroacetone sesqua deutrifum oxide (HFAD) and then diffusion with 300 µL of CBDS.

1H NMR Results: Mole % Composition					
Sample ID	PLA	PGA	PCL		
Foam IA	47.2	12:4	40.5		
Foam IB	12.3	51.3	36.5		
Foam IC	7.7	56.5	35.8		
Foam ID	7.8	56.3	35.8		

[0095] The NMR results indicate that the foam samples have a gradient with respect to composition. The top layer of the foam is high in PLA concentration (47 mole %), whereas the bottom layer of the foam is high in PGA concentration (56 mole %). These results suggest that the PCL/PGA copolymer and the PCL/PLA copolymer have dillerences in their phase separation behaviors during the freezing step and formed a unique compositionally gradient foam.

Example 5

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Transstructural foam

[0096] This example describes the making of a foam that has a compositional and structural gradient and not necessarily a morphological gradient. Such a foam is made from polymer solutions that have been made by physical mixtures of two or more polymers. This example describes a transcompositional foam made from 55/65 PCIL/PLA (as described in Example 9) and 95/5 PLA/PCL (a random copolymer with an IV of 1.8 in HFIP measured as described herein). Note, 95/65 PCIL/PLA (as a soft elastomeric copolymer with 95/5 PLA/PCL is a relatively stiff copolymer, the combination of the two provides a compositional as well as structural gradient. This foam is made using the steps outlined in Example 4 starting from a homogeneous 50/50 physical mixture of a 10% w.f.w. Solution of 95/5 PLA/PCL in 1.4 dioxane. Such a transcompositional foam will provide a good template for tissue junctions such as bone acting en interaction.

Example 6

Cell culture and differentiation data

[0097] Films made from 95.6 PLAPGA, 90/10 PGAPLA, 95/6 PLAPCL, 75/25 PGAPCL and 40/60 PCL/PLA were tested. Tissue culture polythyrene (TCPS) was used as a positive control for all the assays. Before testing, polymer discs were positioned at the bottom of a 24-well ultratow cluster dish and were pre-wetted in govern mode for 20 min. [0098] The 95/6 PLAPGA copolymer used in this example was a random copolymer with an IV of 1.76 as determined in HFIP at 25°C, which is currently used in Parascryl™ suture (Ethicon Inc., Somerville, New Jersey). The 90/10 PGAV PLA copolymer was a random copolymer with an IV of 1.74 as determined in HFIP at 25°C, which is currently used in Polyl™ suture (Ethicon Inc., Somerville, New Jersey). The 95/10 PLAPCL optomer was made as described in Example 91.0 with an IV of 2.1 as determined in HFIP at 25°C. The 75/25 PG/PCL copolymer is a segmented block copolymer with an IV of 1.85 and is described in US Patents, 133.73°9 this copolymer is currently used in Monocryl™ suture (Ethicon Inc., Somerville, New Jersey). The 40/60 PCL/PLA copolymer used in this Example was made as described in Example 9 and had an IV of 1.44.

[0099] Cell attachment and proliferation: Cells were seeded at 40,000 well in 24-well ultralow cluster dishes (Coming) containing the polymers. The ultralow cluster dishes are coated with a layer of hydrogal polymer, which related protein and cell achiesion to the wells. Cell attachment to the biopolymers was determined following 24 his of incubation (N=3 for each polymer). The attached cells were released by typesinization and the number of cells was determined using a hearnecytometr. Cell proliferation was assessed by determining cell counts at days 3 and 6 following seeding.

Differentiation assays:

[O100] Alkaling phosphalase activity. Alkaline phosphalase activity was determined by a colorimetric assay using pnitrophenol phosphale substrate (Sigma 104) and following manufacturers instruction. Briefly, cells were seeded on the films or meshes at a density of 40,000 cellskwell and incubated for 1,6.14, and 21.4 Once cells reached confluence at day 6 they were fed with mineralization medium (growth medium supplemented with 10mM P-gbcerophosphate, 50 ug/mil accorbic acid). Alkaline phosphatase activity was determined in cell homogeniates (0.05%. Tricin X-100) at the above time points. The quantity of protein in cell extracts was determined by micro BCA reagent from Pierce. Primary and ostocolastic cultured on films and meshes were also stained for membrane-bound alkaline phosphatase using a histochemical staining kit (Sigma). For all the films and meshes three samples per arous were tested.

[0101] Osteocalin ELISA: Osteocalcin secreted into the medium by osteoblasts cultured on various films was quantified by ELISA (Osteocalcin ELISA ki), Biomedical Technologies inc, Boston). Aliquots of media from the wells containing the polymer films were hypothilized prior to measurements of this protein by ELISA. Three samples for each polymer were tested and the ELISA was repeated twice.

Von Kossa staining

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[0102] Three samples for each polymer were stained for mineralized tissue using Von Kossa silver nitrate staining.

Expression of alkaline phosphatase and osteocalcin mRNAs

[0103] The expression of alkaline phosphatase and osteocalcin mRNAs in cells was assessed by semi-quantitative reverse transcriptase polymerase chain reaction (RT-PCR) using RNA extracted from cells cultured for 21 d on the films. Seven days after seeding, the culture media was repleased with mineralization media (3 mM ½-glycerophosphate and 50 µg/mi of ascorbic acid were added). The cells were cultured for additional 2 works, for a total period of 3 weeks. Total RNA was executacid from four samples per group using a RNessy mini kit provided by Caigen. The quality and amount of total RNA was measured for each polymer group. Total RNA was everse transcribed to obtain cDNA will are reverse transcriptase reaction (Superscript II, Globo). The cDNAs for osteocation, alkaline phosphatase, and dilyceraldehyde-3-phosphate-dehydrogenase (GAPDH) were amplified using a PCR protocol described previously (Globa) RIL, manufacturors instruction). The primar sequences (Table II) for osteocation, alkaline phosphatase, and GAPDH were obtained using the FASTa program (Genetic Computer Group, Madison, W). Preliminary studies were also conducted to optimize the number of PCR cycles for each primer (Table II), and to determine the range of RNA, which bromide. The gels were photographed under UV light and were evaluated by densitometry for the expression of oster-ceation and additione phosphatase mRNAs relative to GAPDH.

Statistical Anlysis

[0104] Analysis of variance (ANOVA) with Tukey post hoc comparisons was used to assess levels of significance for all the assays.

Table I

Gene	Species	Forward primer	Reverse primer	61
Alkaline phosphatase	Rat	5' ATCGCCTATCAGCTAAT	5'	Size (bp
Osteocalcin	Rat/Human	GCAC 5'CAACCCCAATTGTGA CGAGC	GCAAGAAGAAGCCTTT GGG 5' TGGTGCGATCCATCAC	339
GAPDH	Mouse/ Human/ Rat	5'ACCACAGTCCATGCC ATCAC	AGAG S'TCCACCACCCTGTT GCTGTA	452

Table II

PCR optimization cycles				
Gene	C _{DNA} (μI)	Cycles 25		
Alkaline phosphatase	1			
Osteocalcin	1	35		
GAPDH	1	23		

Results

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[0105] Cell attachment and prollieration on bioresorbable polymens: No observable difference in cell morphology was evident between the various belopyriner films and as compared to TCPS. Cell attachment to the various biopyriner films was equivalent to TCPS following 24 hol incubation. At day 3, cells proliferated well on all films with the exception of 40/60 PCL/PLA, where proliferation was 60% relative to TCPS. Furthermore, 95/5 PLA/PGA and 90/10 PGA/PLA films supported a significantly (p-0.05) higher degree of cell proliferation compared to TCPS and 40/60 PCL/PLA (Fig. 7A).

Differentiation assay:

[0105] Alkaline phosphatase enzyme activity. The profile for alkaline phosphatase activity expressed by osteoblasts cultured on 9575 PLAPGA, 90/10 PGAPILA and 95/5 PLAPGL films was similar to the profile observed on TCPS. Alkaline phosphatase specific activities were significantly (p<0.05) elevated for osteoblasts cultured on 40/60 PCL/ PLA and 75/25 PGAPGL films at days 14 and 21, respectively, compared to other films and TCPS (Fig. 78)

Expression of alkaline phosphatase and osteocalcin mRNA;

[0107] The expression of mRNAs for alkaline phosphalase, osteocalcin, and GAPDH for osteoblasts cultured on the 95/5 PLA_PGA_4060 PLA_PCL_95/5 PLA_PCL_films, and TCPS were evaluated by densitometry. The results are depicted in Fig. 7C. It should be noted that the data in Fig. 7B is at best semi-quantitative. Nevertheless, the data suggests that 40/60 PCL_PLA film supported significantly (p-0.05) higher levels of osteocalcin expression compared to TCPS. The rest of the polymer surfaces were equivalent to TCPS for both osteocalcin and AP mRNAs expression,

Conclusions

[0108] No major differences were observed with respect to cell attachment and proliferation between the different bioresorbable films or meshea tested following 6 days of incubation. Furthermore, the results indicate that differences between these materials were more obvious with respect to their differentiation characteristics. Cells cultured on 40°D0 PCL/PLA film showed enhanced alkaline phosphatase activity and osteocation mRNA expression compared to other films and TCPS following 14 and 21 days of incubation, respectively.

[0109] References that may be referred to for a more complete understanding of this techniques include, M. A. Aronow, L.C. Gerstenledt, T.A. Owen, M.S. Tassinari, G.S. Stein and J.B. Laun. "Factors that promote progressive development of the osteoblast phenotype incultured fetal rat calvaria cells". Journal of Cellular Physiology, 143: 213-221 (1990) and Stein, G. S., Lian, J. B., and Owen, L. A. "Relationship of cell growth to the regulation of tissue-specific gene expression during osteoblast differentiation" ARSEB, 4, 3111-3128 (1990).

Example 7

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In vivo Study of Foam Blend in Swine Dermal Wound Healing Model

[0110] This example describes the results of implanting a 1mm, 0.5mm thickness foam tissue scalfolding in a swine full thickness excisional wound model. The foam tissue scalfold was made from a blend of 40/60 e-caprolactone-co-described as described in Example 3 these polymors were blended together and formed into him and 0.5 mm foams substantially as described in Example 3 (except that the cooling rate was 2.5°C per minute and it was cooled only to 5°C). Scanning electron micrographs of a 0.5mm foam are presented in Figures 9A, 98 and 9C. The two thickness (0.5mm and 1 mm) of toams were then tested in the

wound excisional model with and without PDGF being provided. The resulting four different samples were then evaluated.

[O111] A blinded histologic evaluation was performed on 48 full thickness excisional wounds from four pigs (12 sites per animal) explanted at 8 days following wounding. The assessment was performed on H&E stained sides. During the histologic assessment, the following parameters were ranked evaluated across the specimen set 1) cellular invasion of the matrix qualitative and quantitative assessments 2) infiltration of polymorphonuclear leukocityes (PMNs) into the contact zone (ventral surface) of the matrix, 3) inflammation in the granulation tissue bed below (ventral to) the matrix, 4) reaction of the epidermis to the matrix, and 5) degree of fragmentation of the matrix.

Animal Husbandry:

[0112] The pigs were housed individually in cages (with a minimum floor area of 10-sq. ft.) and given identification. All pigs were assigned an individual animal number. A lag was placed on each individual animal cage listing the animal number, species/farian, surgical dates, surgical technique and duration of the experiment and date of euthanasia. Each animal was clearly marked with an animal number on the base of the neck using a permanent marker.

[0113] The animal rooms were maintained at the range of 40 to 70% R.H. and 15 to 24°C (59.0 to 75.2°F). The animals were led with a standard pig chow once per day, but were fasted overnight prior to any experimental procedure requiring anesthesis. Water was available ad hillium. A daily light clark cycle of 12:12 hours was adopted.

20 Anesthesia:

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[0114] On the initial day of the study, days of evaluation and the day of necropsy, the animals were restrained and anesthetized with either an internuscular injection of Tlettamine HCliplus Zolazepam HCl (Felazole, Fort Dodge Animal Health, Fort Dodge, Iowa 4 mg/ml) and Xylaziae (Rompune, Bager Corporation, Agricutrue Dhission, Animal Health, Shawnee Mission, Kanass, 4mg/ml) or Isoflurane (AErrane® Fort Dodge Animal Health, Fort Dodge, Iowa) inhalatory anesthesia (6% vol.) administered via a nose cone. When the animal was in the surgical suite, it was maintained in Isoflurane (AErrane®) rehalatory anesthesia (6% vol.) administered via a nose cone. Food was available after recovery from each procedure.

30 Preparation of the Surgical Site:

[0115] One day prior to the surgical procedure, body weights were measured and the dorsal region of four pigs were clipped with an electric clipper equipped with a #40 surgical shaving blade. The shaved skin was then re-shaved closely with shaving cream and a razor and then rinsed. The shaved skin and entire animal (excluding the head) was then scrubbed with a surgical scrub brush-sponge with PCMX cleansing solution (Pharmaseal® Scrub Care® Baxter Healthcare Corporation, Pharmaseal Division, Valencia, California) and then with HIBICLENS® chlorhexidine gluconate (available from COE Laboratories, Incorporated, Chicago, Illinois). The animal was wiped dry with a sterile towel. Sterile NU-GAUZE* gauze (Irom Johnson & Johnson Medical Incorporated, Arlington, Texas) was placed over the dorsal surface of each animal and secured with WATERPROOF* tape (available from Johnson & Johnson Medical Incorporated, Arlington, Texas). The entire torso region of the animal was then wrapped with Spandage™ elastic stretch bandage (available from Medi-Tech International Corporation, Brooklyn, New York) to maintain a clean surface overnight. [0116] On the day of surgery, immediately prior to delivering the animal to the surgical suite, the dorsal skin was again scrubbed using a surgical scrub brush-sponge with PCMX cleansing solution (Pharmaseal® Scrub Care®), rinsed and wiped dry using a sterile towel, as performed on the previous day. The animals were placed prone on the surgical table and wiped with 70% alcohol and dried with sterile gauze. Using a sterile surgical marker (available from Codman® a division of Johnson & Johnson Professional Incorporated, Raynham, Massachusetts) and an acetate template, marks were made on the dorsal skin according to the desired placement of each full-thickness wound,

Surgical Procedure:

[O117] Following anesthesia, under sterile conditions, twelve (12) full-thickness excisions (1.5 x 1.5 cm) per animal were made in two rows parallel to the spiral column on the left and right dorsal regions using a scaled blade. A pair of ecisions and/or scatpol blade was used to a did in the romoval of skin and subcutaineous tissue. Blooding was controlled by use of a sponge tamponade. Sufficient space was fit between wounds to avoid wound-to-wound interference. The excised tissue was measured for thickness using a digital calipits.

Application of the Treatment and Dressing:

[0118] Each wound was submitted to a prepared, coded treatment regimen (study participants were blinded to all treatments). The primary dressing consisting of the sterile individual test article (1,5 x 1,5 cm coaked in sterile satins for 24 hours) was placed into the wound deficit in a predetermined scheme. The secondary dressing, a non-arberent, saline scaked, square of RELEASE' dressing (manufactured by Johnson & Johnson Medical Incorporated, Affington, Texas) was placed on top of the test article. A layer of BIOCLUSIVE' dressing idevaliable from Johnson & Johnson Medical Incorporated, Affington, Texas) was sealed over the wounds to keep the wound moist and the dressing in place. Strips of Reston** (8M Medical-Surgical Division, St. Paul, Minnescola polypurethera self-arbering foam verp placed between the wounds to avoid cross-contamination due to wound fulful leakage, and to protect the wounds from dranage and the dressing from displacement. A layer of NU-GAUZE' gauze was placed on top of the BIOCLUSIVE' dressing and Reston** foam, and was secured with WATERPROOF' tape to protect the dressings. The animals were then dressed with Spandage** elastic not to he keep the versings in place.

[0119] The secondary dressings were removed and replaced daily with a fresh piece of saline scaked RELEASE* secondary dressing. The primary dressings (test articles) were not disturbed unless the unit was displaced or pushed out of the wound delicit.

POST-OPERATIVE CARE AND CLINICAL OBSERVATIONS:

[0120] After performing the procedures under anesthesia, the animals were returned to their cages and allowed to recover. The animals were given analysatics (buprenorphine hydrochloride [Buprenex Injectable, 0.01 mg/kg, in] sold by Rockitt & Colman Products, hult, Englandy immediately post-surgery and the following day. After recovering from anesthesia, the pigs were observed for behavioral signs of discomfort or pain. No signs of pain were observed.

[0121] Each pig was observed twice daily after the day of surgery to determine its health status on the basis of general attitude and appearance, food consumption, focal and urinary excretion and presence of abnormal discharges.

EUTHANASIA:

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[0122] At the end of the study (6 days post-wounding), each animal was euthanized under anesthesia, with an iniravenous injection of (1 ml/10 pounds body weight) Socumb™ pentobarbital sodium and pherytoin sodium euthanasia solution (sold by The Butler Company, Columbus, Chio) via the marginal ear vein. Following euthanasia, the animals were observed to ensure that respiratory function had ceased and there was no palpable cardiac function. A stethoscope facilitated the assessment for the lack of cardiale function.

35 TISSUE HARVESTING:

[0123] Immediately following euthanasia, each wound, together with the underlying fat and a small portion of surrounding skin was excised. The tissue was placed in 10% neutral buffered formalin.

#0 EVALUATIONS:

Visual Wound Assessment:

[0124] General observations were recorded for days 1-3, including displacement, wound reaction and physical characteristics of the scatfold. Detailed clinical evaluations were performed on days 4 - 8 post-wounding. Assessments were recorded as to the presence/absence (yes = 1/no = 0) and/or degree (given a score) of the following parameters:

<u>Dressing Conditions</u>: air exposed, displacement of test article, channeling, communication and moisture content of the RELEASE* secondary dressing(scored as: 4=moist, 3=moist/dry, 2=dry/moist, 1=dry).

Wound Bed Conditions; moisture content of test article (scored as: 4=moist, 3=moistdry, 2=dry/moist, 1=dry), inflammation (scored as: 3=severe, 2=moderate, 1= slight, 0=none), reinjury (scored as: 3=severe, 2=moderate, 1= slight, 0=none), clots, folliculitis, infaction, level of test article (scored as: 4=super raised, 3=raised, 2=severe, 1=doprossed), fibrin (scored as: 3=severe, 2=modorate, 1= slight, 0=none), and drythoma. Color of the tost article was also observed.

TISSUE PROCESSING:

[0125] Excised tissue samples were taken at day eight. The entire wound was harvested and placed into 10% neutral

buffered formalin. The tissue was prepared for frozen sections. The tissue was trimmed and mounted onto the object holder with Tissue-Tek® OCT Compound (sold by Sakura Finetechnical Company, Limited, Tokyo, Japan) and quickty frozen. The specimens were sectioned on the cryostal at 10 µm and stained with a frozen H&E stain.

5 HISTOLOGICAL ASSESSMENTS (Day 8 post-wounding):

[0126] Histological evaluations for granulation tissue (area and length) and epithelialization were assessed using H&E stained specimens using a magnification of 20-40X. Granulation tissue height was determined by dividing the area by the length.

[0127] Histopathological evaluation of the tissue samples was assessed using the H&E stained specimens, they were first assessed under 100x to 400x magnification.

RESULTS

[0128] There was cellular invasion into the interstices of the matrix in the majority of all test sites. In the majority of sites this invasion was true its sue ingrowth comprised of varying populations of libroblasts, macrophages, macrophage giant cells, and endothelial-like cells, there also appeared to be capiliary formation. Significant formation of dense librous connective itssue bayer dorsal to the matrices essentially embedding the matrices in the tissue, was seen at several sites for the 0.5mm forams with and without POEr. The 1 mm matrices were either at the surface of the tissue bed or sloughed. Macrophage giant cell formation seemed to be greater in the 0.5mm versus the tirm loam scallots, in sites where the 1 mm foam scallots provided the site of the 1 mm foam scallots.

[0129] Complete incorporation of the matrix into the granulation tissue bed was seen at several sites for the 0.5mm loam scaffoldings. Figures 10 and 11 illustrate the incorporation of these matrices into the granulation tissue bed. Figure 10 is a dark filed 40X pictomicrograph of a trichrome stained tissue sample.

[0130] Figure 11 is a 100X composite photomicrograph of a trichrome stained sample illustrating the cellular invasion of a foam containing PIOSF. Complete incorporation of the matrices into the granulation tissue bed is evident in both pictures. The dense fibrous tissue above the foam scaffolding is evident in both pictures. These results indicate the 0.5mm foams will provide a suitable substrate for the growth of epidermal tissue.

Example 8

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Synthesis of a Random Poly(ε-caprolactone-co-glycolide)

35 [0131] A random copolymer of e-caprolactone-glycolide with a 35,65 molar composition was synthesized by ring opening polymerization reaction. The method of synthesis was essentially the method described in U.S. Patent 5,468,255 in Example 6 (which is hereby incorporated herein by reference). The amount of delthylene glycol initiator added was adjusted to 1.15 mmole/mole of monomer to obtain the following characteristics of the dride polymer. The inherent viscosity (I.V.) of the copolymer was 1.59 dL/g in hexafluoroisopropanol at 25°C. The molar ratio of PCL/PGA was found to be 35.5/64.5 by protion NMR with about 0.5% residual monomer. The glass transition (Tg) and the melting points (Tm) of the copolymer were found to be -1°C, 60°C and 126°C respectively. by DSC.

Example 9

45 Synthesis of 40:60 Poly(ε-caprolactone-co-L-lactide) by Sequential Addition

[0132] In the glove box, 100 µL (33 pmol) of a 0.33 M stannous octoate solution in toluene, 115 µL (1.2 mmol) of disthylene glycol, 24.6 grams (170 mmol) of Liscitide, and 45.7 grams (400 mmol) of Leoprolactione were transferred into a silentized, filame dried, two neck, 250 mL round bottom flask equipped with a stainless steel mechanical stirrer and a nitrogen gase blanket. The reaction flask was placed in an oil bath already set at 190°C and held there. Meanwhite, in the glove box, 62.0 grams (400 mmol) Liscitide were transferred into a flame dried, pressure equalizing addition funnet. The furnel was wrapped with heat tape and attached to the second neck of the reaction flask. After 6 hours at 190°C, the molton Liscitide was added to the reaction flask over 5 minutes. The reaction was continued overnight for a total reaction time of 24 hours at 190°C. The reaction was allowed to cool to room temperature overnight. The copolymer was isolated from the reaction flask by freezing in liquid nitrogen and breaking the glass. Any remaining glass fragments were removed from the copolymer using a bench grinder. The copolymer was again frozen with liquid nitrogen and breaking the glass. Any remaining plass fragments were removed from the copolymer using a bench grinder. The copolymer was again frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with liquid nitrogen and breaking the glass plan frozen with

co-Lacifide) were added to a tared aluminum pan and then devolitilized under vacuum at 110°C for 54 hours, 98.7 grams (95.7% by weight) of copolymer were recovered after devolitilization. The inherent viscosity was measured and found to be 1.1 d/Lg in ChCl₃ at 26°C (c = 0.1 g/dL). FTIR (cast film from ChCl₃ onto KBr window, cm**): 2993, 2944, 2868, 1759, 1456, 1393, 1392, 1194, 1132, 1094, 870, and 756. ¹H NMF, (400MHz, HFA/Denzene, ppm): 8 1.25. 2 broad lines; 40, 1.55, 2 lines; 22.5, 5 lines; 2.25, 5 lines; 2.35, 5 lines; 2.35, 5 lines; 2.35, 6 li

10 Example 10

Synthesis of 95/5 PLA/PCL Copolymer

[0133] In the glove box, 170 µL (18 mmol) of diethylene glycol, 350 µL (115 pmol) of a 0.33 M stannous octoate solution in toluene, 17.1 grams (150 mmol) of e-caprolactone, and 410.4 grams (2.85 mol) of L-lactide were placed into a silanized, flame dired, 1000 mL round bottom equipped with a stanless steel mechanical strere and vacuum take off connector in order to maintain a dry nitrogen gas blanket. The reaction flask was placed in an oil bath already heated to 185°C and then held there for 3 hours. The flask was removed from the oil bath and allowed to cool down to room temperature. The polymer was isolated by wrapping the flask with aluminum foil, freezing it in fliquid nitrogen, and then grinding away any adhered glass to the polymer. The copolymer was then ground in a Wiley mill. The ground polymer was vacuum dried at 60°C for 24 hours. 30°2 grams of copolymer were collected. The inherent viscosity was 2.1 d./g in chloroform [25°C, c=0.1 g/d.]. The copolymer composition was measured by proton NMF spectroscopy and found to be 97.2 mole percent PLA and 2.8 mole percent PLA net 75.

Claims

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- A biocompatible gradient foam having a first location and a second location wherein the biocompatible gradient foam has a substantially continuous transition in at least one characteristic selected from composition, stiffness, flexibility, bioasbsorption rate and pore architecture from the first location to the second location of said biocompatible gradient foam.
- 2. A biocompatible foam having a first surface and a second surface with interconnecting pores and channels.
- 35 3. The foam of claim 1 or claim 2 which is bioabsorbable.
 - 4. The foam of any one of claims 1 to 3 which is made from a bioabsorbable aliphatic polyester, a poly(amino acid), a copol/(ether-ester), a polyalylene oxalate, a polyamide, a poly/(minocarbonate), a polyorthoester, a polyoxaester, a polyamidoester, a polyoxaester containing amine groups, a poly(anhydride), a polyphosphazene, a biopolymer or a blend thereof.
 - 5. The foam of claim 4 wherein additionally present as a constituent of the foam is a second aliphatic polyester.
 - The foam of any one of claims 1 to 5 wherein also present in the foam is a therapeutic agent.
 - The foam of claim 6 the therapeutic agent is an antiinfective, a hormone, an analgesic, an anti-inflammatory agent, a growth factor, a chernotherapeutic agent, an anti-rejection agent, a prostaglandin, an RDG peptide or a combination thereof.
- 50 8. The foam of any one of claims 1 to 7 wherein the biocompatible gradient foam is filled with a biocompatible material selected from bioabsorbable synthetic polymers, biocompatible, bioabsorbable biopolymers, biocompatible ceramic materials and combinations thereof.
- The foam of claim 2 or any claim dependent thereon having interconnected pores formed from a composition containing in the range of from 30 weight percent to 99 weight percent e-caprolactione repeating units.
 - 10. The foam of claim 9 having a first location and a second location wherein the biocompatible foam has a substantially continuous transition in at least one characteristic selected from composition, stiffness, flexibility, bioabsorption

rate and pore architecture from the first location to the second location of said biocompatible foam.

- 11. The foam of claim 9 or claim 10 wherein the interconnecting pores have a pore size in the range from 10 μm to 200 μm.
- 12. The foam of any one of claims 9 to 11 wherein the biocompatible foam has a porosity of in the range of from 20 to 98 percent.
- 13. The foam of any one of claims 9 to 12 wherein the biocompatible foam has channels.

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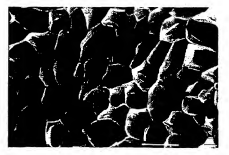
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- 14. The foam of claim 13 wherein the channels have an average length of at least 200 μm .
- 15. The foam of any one of claims 1 to 14 which is formed with an insert within the biocompatible gradient foam.
- 5 16. The foam of any one of claims 1 to 15 for use in the repair or regeneration of tissue.

FIG. 1

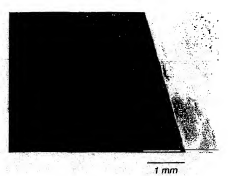


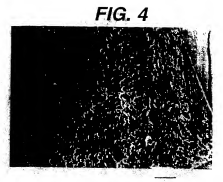
100 µm

FIG. 2

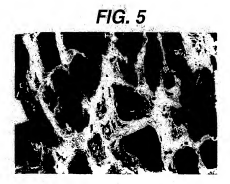


FIG. 3





1 mm



100 μm

FIG. 6



100 μm

FIG. 7A

% Cell Proliferation on Bioresorbable Films at Day 3 and Day 6 relative to TCPS

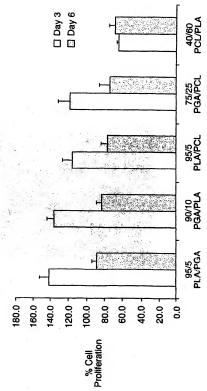


FIG. 7B

Alkaline Phosphatase Specific Activity on Bioresorbable films

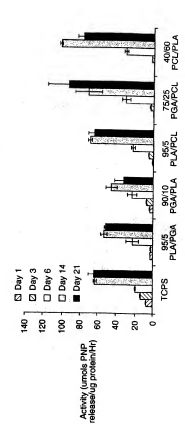
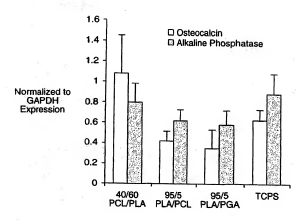


FIG. 7C



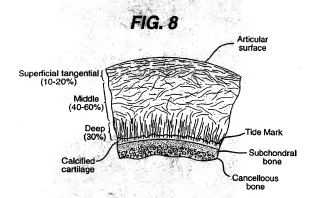
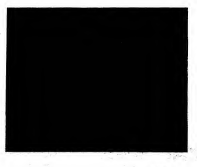
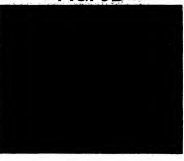


FIG. 9A



150 µm

FIG. 9B

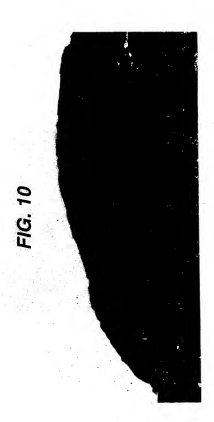


150 µm

FIG. 9C



100 um









EUROPEAN SEARCH REPORT

Application Number

EP 00 30 5501

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